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EFFECT OF TWO-STAGE AGING ON MICROSTRUCTURE OF 7075 ALUMINUM ALLOYS



RE- 627

Final Report



April 1981

by

John M. Papazian

Prepared Under Contract N00019-79-C-0285



for

Department of the Navy Naval Air Systems Command Washington, D.C. 10351

by

Research Department Grumman Aerospace Corporation Bethpage, New York 11714

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Differential scanning calorime (TEM) were used to establish the r	uny (Dob) and trans Microstauctural of	smission electron microscopy facts of various multistage
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For two-stage aging treatments in which the final aging was for 8 h at 150°C, pre-aging for 6 h at 90°C or equivalent treatments at 120°C. This is in agreement with the Pashley et al model of two-state aging. For two-stage treatments where the first aging treatment is 4 h at 950C, increasing the final aging temperature from 150 to 155 or 160°C resulted in a significant increase in the amount of \mathfrak{n}' in the microstructure; variations in the pre-aging treatment has much less of an effect. It was found that the RRA heat treatment converts the T6, predominately G.P. zone microstructure, to a more stable T7 type of microstructure; but the RRA microstructure consists of smaller precipitates and is more heterogeneous than the T7. The major differences between the T6, two-stage T6, RRA and T73 microstructures appear to be in the relative proportions of G.P. zones, η' and η . The percentage of G.P. zones is greatest in T6, least in T73 and RRA, and intermediate in the two-stage T6. Based on these limited observations, the stress corrosion susceptibility of 7075 appears to be more influenced by the matrix microstructure than by the grain boundary precipitate size and spacing.

ABSTRACT

Differential scanning calorimetry (DSC) and transmission electron microscopy (TEM) were used to establish the microstructural effects of various multistage aging treatments on aluminum alloy 7075. DSC was used to characterize the matrix microstructure while TEM was used to characterize the grain boundary precipitates. The aging treatments were in five categories: isochronal pre-aging, isothermal pre-aging, final aging at 150°C, commercial type two-stage aging treatments, and retrogression and re-aging (RRA) treatments.

For two-stage aging treatments in which the final aging was for 8 h at 150°C, pre-aging for 6 h at 90°C produced a slightly finer and harder microstructure than longer treatments at 90°C or equivalent treatments at 120°C. This is in agreement with the Pashley et al model of two-stage aging. For two-stage treatments where the first aging treatment is 4 h at 95°C, increasing the final aging temperature from 150 to 155 or 160°C resulted in a significant increase in the amount of n' in the microstructure; variations in the pre-aging treatment had much less of an effect. It was found that the RRA heat treatment converts the T6, predominately G.P. zone microstructure, to a more stable T7 type of microstructure; but the RRA microstructure consists of smaller precipitates and is more heterogeneous than the T7. The major differences between the T6, two-stage T6, RRA and T73 microstructures appear to be in the relative proportions of G.P. zones, η' and η . The percentage of G.P. zones is greatest in T6, least in T73 and RRA, and intermediate in the two-stage 16. Based on these limited observations, the stress corrosion susceptibility of 7075 appears to be more influenced by the matrix microstructure than by the grain boundary precipitate size and spacing.

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INTRODUCTION

1.1 BACKGROUND

The poor stress corrosion resistance of aluminum alloy 7075 has led to numerous attempts to circumvent this problem. Newer alloys in the Al-Zn-Mg family have been developed, and overaging techniques for 7075 have been devised. Despite significant activity in this area, an entirely satisfactory solution has not yet been obtained. Two recent attempts at property improvement for 7075 involve two-stage aging treatments (Ref 1) and a patented, retrogression and re-aging (RRA) treatment (Ref 2). These treatments aim at producing precipitate microstructures in 7075 which have strength levels equal to those of the commonly used T6 temper but a stress corrosion resistance which is equal to or greater than that of the T73 temper. The two-stage aging treatment of 6 h at 105°C followed by 8 h at 150°C has been reported to increase stress corrosion life by serveral hundred percent while conserving the same tensile properties as material aged for the conventional 24 h at 120°C (Ref 1). Likewise, the RRA treatment can increase the stress corrosion threshold stress up to three times the normal value in 7075-T6 (Ref 2).

The objective of this work was to quantitatively characterize the precipitate microstructures of 7075 after systematic variations in two-stage aging treatments in order to establish the manner in which these precipitates depend upon the aging parameters. The selected aging heat treatments centered around the two-stage aging treatment reported by Di Russo, et el (Ref 1 and 3) and the various two-stage commercial treatments for T6, T73 and T76 (Ref 4). In addition, a sample of 7075-T651 which had received the patented RRA treatment was evaluated. The matrix microstructural evaluations were performed using differential scanning calorimetry (DSC). In previous studies, DSC has been shown to be of use for rapid characterization of the microstructures of 7000 series aluminum alloys (Ref 5 and 6). Transmission electron microscopy (TEM) was employed to determine the nature of the grain boundary precipitate microstructures, and electrical conductivity (EC) and hardness were used to further characterize the material.

1.2 PREVIOUS WORK

The sequence of precipitate phases observed during the aging of the A1-Zn-Mg alloys has been extensively studied (Ref 7 - 10). The aging sequence is currently thought to be (Ref 9 and 11):

supersaturated $\alpha \rightarrow G.P.$ zones $\rightarrow \eta "\rightarrow \eta "\rightarrow \eta$.

The G.P. Zones are spherical and have a f.c.c. structure which is fully coherent with the matrix (Ref 7). The η'' phase is usually described as spherical and coherent, but with an ordered G.P. zone or hexagonal structure (Ref 8, 9 and 11). The η' phase is a partially coherent, plate like precipitate on {111} with a hexagonal structure (Ref 8 and 12). The stable η , MgZn₂, phase also has a hexagonal structure and can transform to a ternary cubic T phase after prolonged aging at high temperatures (Ref 8).

The temperatures at which the different phases form vary with the exact composition of the alloy. Most of the microstructural work has been performed on pure alloys. In general, G.P. zones are the first phase to form at low aging temperatures, T \leq 75°C. When it is detected, η ", is observed at intermediate aging temperatures (75 to 150°C) or after long term aging at low temperatures. It is not always included as part of the aging sequence (Ref 7 and 10). The η' phase is also observed in this same temperature range. The η phase generally occurs at higher temperatures; but the work of Gjønnes and Simensen (Ref 8) and DeArdo and Simensen (Ref 13) suggests that η' and η can both be present after aging at 100°C.

The microstructure of Al-Zn-Mg alloys after two-stage aging treatments has been studied using TEM by Lorimer and Nicholson (Ref 14) and by DeArdo and Simenson (Ref 13). The results of these studies show finer precipitate distributions after two-stage aging and have been interpreted on the basis of theories of two-stage aging developed by Lorimer and Nicholson (Ref 14) and by Pashley, et al (Ref 15). The Lorimer and Nicholson theory postulates that when the low temperature aging treatment is performed helow Tg.p., the highest temperature for G.P. zone precipitation, the G.P. zones thus formed whose diameters exceed a critical value will survive the up-quench to the second aging temperature and provide the nucleation sites for n'. The Pashley et al model is similar except that the precipitates formed at the higher temperatures are thought to nucleate on clusters that are inherited from the low temperature aging treatment. The clusters need not be G.P. zones and may

have the structure of the final precipitate. The Pashley et. al. model was thought by DeArdo and Siemensen (Ref 13) to provide the best explanation for their results, particularly because they thought η' and η were both present at the lower aging temperatures.

* *

In addition to the work mentioned above, a considerable amount of work has been done on the aging of pure Al-Zn-Mg alloys using DSC. This work has been performed by groups working in Japan, France and Austria, and Hungary. The Japanese workers have studied the aging of Al-Zn-Mg (Ref 16), two-step aging of Al-Zn-Mg (Ref 17) and prolonged aging of Al-Zn-Mg (Ref 18). In all of these studies, the alloy used was a pure pseudo-binary Al-6 wt% MgZn₂ with a composition of 0.94 wt% Mg, 5.06 wt% Zn. The calcrimetric experiments reported in these papers were very well done and several of the general observations appear to be applicable to 7075. However, since the alloy used was a pure material whose composition was nearer to that of a medium strength Al-Zn-Mg alloy such as 7011, most of the detailed observations are not applicable. In particular, Hirano showed that the structure produced during aging at 130°C was strongly influenced by a pre-aging treatment at low temperature. This does not seem to be true for 7075 aged at 120°C, as will be shown later. In addition, Hirano's assignment of specific reaction peaks to particular dissolution reactions was not confirmed by TEM, and some choices appear to be questionable.

The French and Austrian workers have also performed extensive calorimetric investigations of A1-2n-Mg alloys (Ref 19). Once again, they have studied pure alloys whose compositions are very nearly 5 wt% Zn, 1 wt% Mg. As with the Japanese studies, many of their detailed observations do not appear to be applicable to 7075. The major differences appear to be shifts in the temperature ranges in which the various transition phases are observed and dissimilar nucleation behavior for G.P. zones. In an early study of 7039, 7075 and a pure A1-Zn-Mg alloy, Thompson (Ref 20) has documented some of these dissimilarities.

The Hungarian workers also used pure alloys, but a variety of compositions were chosen (Ref 21-23). None of their alloys contained copper or chromium, but they documented the presence of the η'' (or G.P. II) phase. They also arrived at a dissolution enthalpy of 2.9 J/g for one volume percent G.P. zones in a variety of Al-Zn-Mg alloys.

EXPERIMENTAL PROCEDURES

The starting material for this investigation was a 6 in. long piece of 1 in. thick by 48 in. wide plate of 7075-T651 which was provided by the Naval Air Systems Command. Samples were machined from this material by first bandsawing 1 in. x 48 in. x 0.1 in. thick slabs from the plate and then milling and cutting to produce 1 in. x 2 in. x 0.063 in. sample coupons. Thus, the 1 in. dimension was parallel to the short transverse direction of the plate, the 2 in. dimension was parallel to the long transverse direction and the 0.063 in. parallel to the longitudinal direction. These coupons were solution heat treated for 2 h at 482°C in flowing argon and drop quenched into room temperature water. They were then transferred to liquid nitrogen and stored at 77 K. Subsequent aging of the samples was performed in silicone oil baths controlled to \pm 1°C.

After aging, the electrical conductivity and hardness of the samples were measured using standard techniques. DSC analyses were performed on hand punched, 7/32 in. diameter, discs using a DuPont 990 Thermal Analyser with a plug-in DSC cell. At least three DSC runs were made for each sample in order to establish reproducibility. All of the DSC runs started at room temperature, ended at 530°C and were made with a constant heating rate of 10°C/min. A pure aluminum disc of approximately equal mass was used as the reference. The thermal analyser was connected to a mini-computer with a suitable interface, and the data from each run were continuously stored. After a run was completed, the data were converted to heat capacity vs. temperature using a previously established calibration for the DSC cell. Subsequently, a linear baseline was subtracted from the data. This baseline represents the temperature dependent heat capacity of the aluminum rich solid solution and existing precipitates, and its value was in agreement with the Neumann-Kopp rule. The remainder, the differential heat capacity ($\Delta C_{\rm D}$), represents the heat associated with solid state reactions which occurred during the DSC run. The reaction peaks in the ΔC_{p} vs T curves can be characterized by a reaction enthalpy, ΔH_r , the area of the peak, which is proportional to the molar heat of dissolution or formation of the precipitate and its concentration; and by T_{r} , the peak temperature, which is the temperature of maximum reaction rate.

This technique and its application to microstructual characterization of 7000 series aluminum alloys has been described in greater detail elsewhere (Ref 5 and 6).

TEM specimens were prepared from 3 mm (0.14 in.) discs that were punched from a section of the original sample that had been hand ground to approximately 0.25 mm (0.01 in.) thickness. Final thinning was performed in a twin jet electropolisher using a one part nitric acid, two parts methanol electrolyte at -20°C. The thin foils were examined in a Phillips EM300 equipped with a goniometer stage. In order to determine the characteristics of the grain boundary precipitates, the thin foil was first rotated such that a grain boundary was aligned with the tilt axis, and then tilted such that the boundary plane was vertical (parallel to the incident beam). The foil was then tilted again by approximately 30 to 40° so that the precipitates in the grain boundary were visible, and the boundary was photographed. Since the angle between the grain boundary and the incident beam was known, the grain boundary area could be accurately calculated. The sizes and densities of the grain boundary precipitates were then measured from the electron micrographs as follows.

The grain boundary precipitates were generally small, circular platelets lying in the plane of the grain boundary, thus the parameters measured were: N, the total number of precipitates; N_A , the number of precipitates per unit area of grain boundary; $\bar{\lambda}$, the average length of the precipitate particle along the grain boundary; and A_f , the area fraction of the grain boundary covered by precipitates. The length of the precipitate along the grain boundary was measured since this direction was the tilt axis of the foil and was therefore not distored by projection. The area fraction was calculated by assuming that the precipitate platelets were disc shaped and had an average diameter equal to $\bar{\lambda}$.

A sample of the original T651 plate that had been RRA treated by Lockheed-California was also analysed. The sample was approximately 0.5 in. x 1 in. x 6 in. and had been taken from the section of the plate labeled 6B3. Specimens for DSC analysis were taken from the face of this sample and from the mid-thickness by slicing with a water cooled, diamond impregnated, crystal cutting saw. DSC was the only technique used to evaluate this material.

RESULTS

The aging treatments used in this study can be arranged into five groups for comparison purposes: isochronal pre-aging treatments, isothermal pre-aging treatment, final aging at 150°C, commercial type two-stage aging treatments and the retrogression and re-aging treatment. Results from the DSC analyses of the microstructures of these five groups of treatments will be presented in sequence. These will then be followed by the results of the TEM study of grain boundary microstructures.

3.1 ISOCHRONAL PRE-AGING TREATMENTS

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Thermograms from samples that were aged for 4 h at 90, 95, and 100°C are shown in Fig. 1. All of the DSC measurements in this study ran from room temperature to 530°C, but in Fig. 1 only the interval between 50 and 300°C is shown. The higher temperature portions of the thermograms have been omitted from the figures because they show only the dissolution of the stable, η , phase and they were identical for all of the aging treatments used. Figure 1 shows a small, endothermic, dissolution peak near 150°C (Peak I) and a large, exothermic, precipitation peak near 225° (Peak II). Aging for 4 i. at 90 or 95°C resulted in similar microstructures, but 4 h at 100°C resulted in a decreased ΔH_r and an increased T_r for both dissolution and precipitation. These changes are tabulated in Table 1 which also lists the hardness and electrical conductivity of these samples. Both the hardness and conductivity of these samples increased as the aging temperature increased. Aging for 4 h at 100°C caused the hardness to increase significantly more than aging at 90 or 95°C. Note that the hardness values measured throughout this work tended to be slightly lower than expected for commercial practice. This is thought to be due to the use of small samples and the absence of stretching or stress relieving treatments.

The results of aging for 6 h at 75, 85, 90, 95, 100, 105 or 120°C are shown in Fig. 2. The trends of these data are similar to those shown in Fig. 1. As the aging temperature increased, ΔH_{Γ} for both peaks decreased. Simultaneously, T_{Γ} for both peaks increased. Peak II, the formation peak, has two maxima (Peaks IIa and IIb) after the lower temperature aging treat-

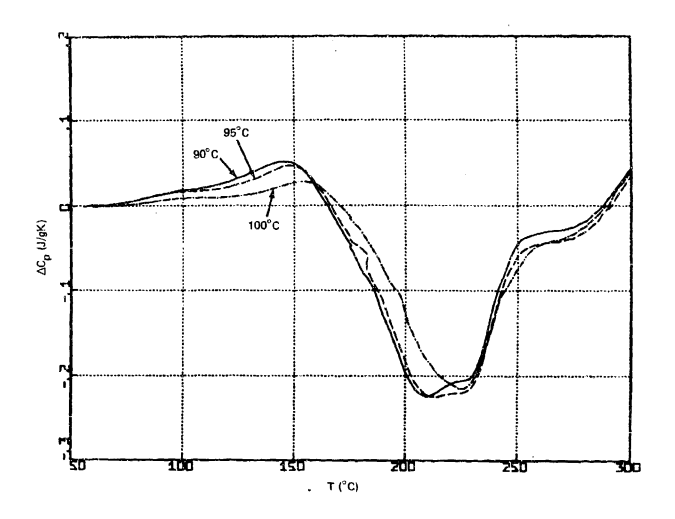


Fig. 1. DSC Traces From Solution Treated 7075 After Aging for 4 Hours at 90, 95 and 100°C.

Table 1. DSC Characteristics After Isochronal Pre-Aging Treatments

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Aging				Peak I		l Pe	ak IIa	Peak 11b	
t (h)	T (°C)	Hardness (R _B)	EC (% IACS)	Τ _r (°C)	ΔH _r (J/g)	т _г (°С)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g
4	90	60±10	30.0	147	2.4±,1	211±2	-13.2±.4		
4	95	64±4	30,1	148	2,3±.2	215±3	-12.8±.9		
4	100	77±3	30.2	154	1.3±.1	224	-11.5±.1		
6	75	66±8	29,8	137±1	4.2±.5	208	·8.4±.2	228	-4.9±.6
6	85	72±2	30,0	149	3.0±.1	213±2	-12.7±.4		
6	90	€5±2	30.8	146	3.1±.3	211±2	7.4±.5	230	4,4±.8
6	95	67±7	30.5	154	1.7±.4	225	-11.7±.3		
6	100	76±4	30.6	155	1.2±.4	225	-11.9±.5		
6	105	71±4	31.0	159	1.45±.3	225	-9.9±,4		
6	120	69±4	31.5	188	.31±.3	225	-7.5±.4		
8	95	76±4	30.3	156±2	2.0±,4	225	·11.4±.3		
8	100	70±4	31.0	159±1	.8±,2	226	-11,2±,2		
8	105	76±4	31.0	180	.9±.7	226±2	-9,3±,3		
8	110	76±3	31.0	174±7	.7±.5	227±1	-9,2±,3		

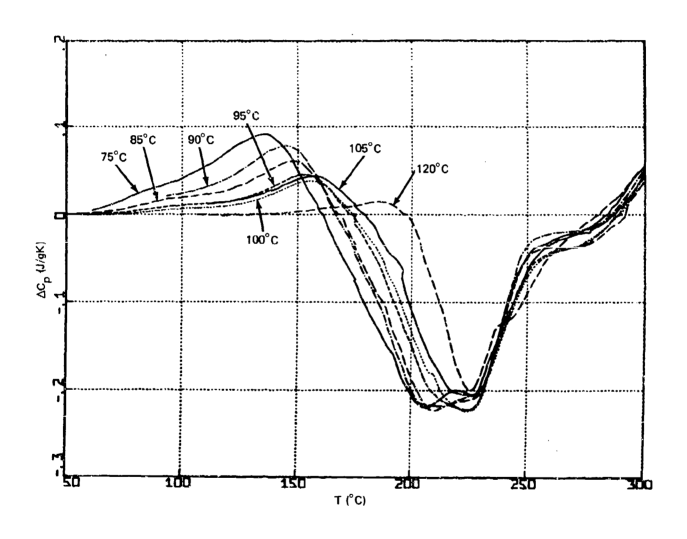


Fig. 2. DSC Traces From Solution Treated 7075 After Aging for 6 Hours at 75, 85, 90, 95, 100, 105 and 120°C.

ments, but evolves into a single peak after aging at the higher temperatures. (Note that the extrema appear as minima in the ΔC_p vs T plots but they are called maxima because they represent a maximum exothermic effect.) Values of the peak temperatures and enthalpies are given in Table 1, along with the hardness and conductivity. The conductivities of these samples increase as the aging temperature increases, but the hardness values have too much scatter to show a clear trend.

Results from the final series of isochronally pre-aged samples are shown in Fig. 3. Aging for 8 h at 95, 100, 105 and 110°C resulted in microstructural trends which are similar to the 4 and 6 h series. Peaks I and II both moved to slightly higher temperatures as the aging temperature increased, and their respective enthalpies decreased. The tabulated results in Table 1 show that the conductivities increased as the aging temperature increased but the hardness values did not change significantly.

3.2 ISOTHERMAL PRE-AGING TREATMENTS

The results of aging for 4, 6, 24, and 96 h at 90°C are shown in Fig. 4. Longer aging times resulted in an increased ΔH_r and T_r for Peak I. Concurrently, ΔH_r for Peak II decreased and T_r for Peak II increased. During these aging treatments, Peak II also evolved from a composite, double maxima, shape into a simpler peak with only one maximum. This change in shape of Peak II was also seen during isochronal aging, as previously shown in Fig. 1 and 2. Table 2 lists the characteristics of these samples after heat treatment. The hardness and conductivity both increased significantly as the aging time increased.

The results of aging for 4, 6, and 8 h at 95°C are shown in Fig. 5. Increased time at the aging temperature resulted in a small increase in T_Γ for Peak I and an evolution of the shape of Peak II. These changes were similar to those noted above for aging at 90°C but they were not as pronounced because of the smaller range of aging times studied. The results tabulated in Table 2 show the same general trends in hardness and conductivity as observed for aging at 90°C.

Figure 6 shows the results of aging for 4, 6, and 8 h at 100°L. As the aging time increased, there was a small decrease in the ΔH_{Γ} of Peak I

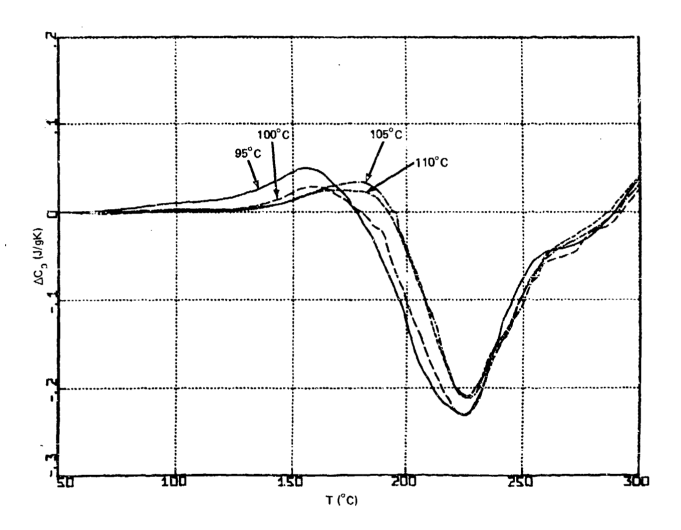


Fig. 3. DSC Traces From Solution Treated 7075 After Aging for 8 Hours at 95, 100, 105 and 110°C.

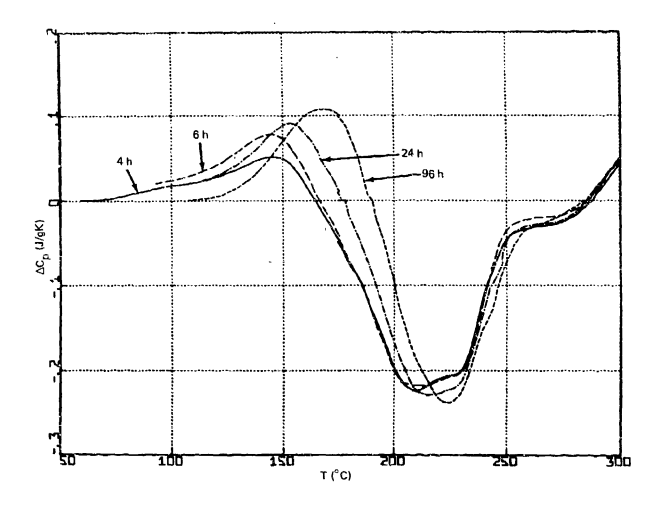


Fig. 4. DSC Traces From Solution Treated 7075 After Aging for 4, 6, 24 and 96 Hours at 90°C.

Table 2. DSC Characteristics After Isothermal Pre-Aging Treatments

Aging				P	eak I	Pe	ak IIa	Peak IIb	
t (h)	T (°C)	Hardness (R _B)	EC (% IACS)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)
4	90	60±10	30.0	147±1	2.4±,1	211±2	-13.2±.4		
6	90	65±2	30,8	146±1	3.1±.3	211±2	-7.4±.5	230±1	4.4±.8
24	90	72±2	30.3	154±1	3.5±.1	218±5	-11.5±.8		
96	90	80±3	30.9	171±2	4.2±.2	224±2	·10.0±.5		
4	95	64±4	30.1	148±1	2.3±.2	215±3	-12.8±.9		
6	95	67±7	30.5	154±1	1.7±.4	225±1	-11.7±,3		
8	95	76±4	30.3	156±2	2.0±.4	225±1	·11.4±.3		
4	100	77±3	30,2	154±1	1.26±.1	224±1	-11.5±.1		
6	100	76±4	30.6	155±1	1.2±.4	225±1	11.9±.5		
8	100	70±4	31.0	159±1	.8±.2	226±1	-11.2±.2		
6	120	69±4	31.5	188±1	.3±.3	225±1	-7.5±.4		1
24	120	87±1	32.3	192±2	4,7±,3	227±1	-3.7±.3	246±3	-3.1±.1
96	120	86±1	33.0	192±2	5,9±,2	227±1	-2.6±.2	252±1	-2.7±.2

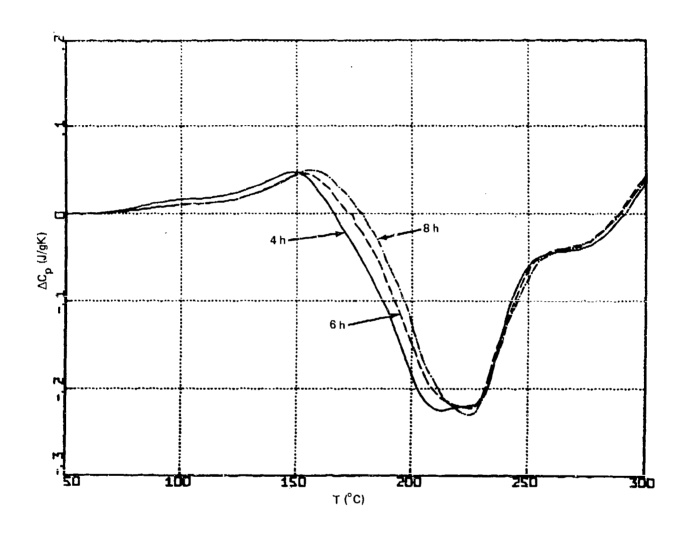


Fig. 5. DSC Traces From Solution Treated 7075 After Aging for 4, 6 and 8 Hours at 95°C.

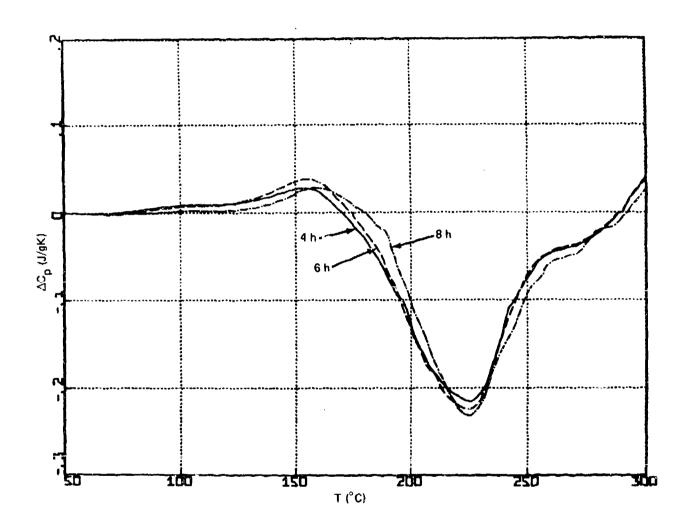


Fig. 6. DSC Traces From Solution Treated 7075 After Aging for 4, 6 and 8 Hours at 100°C.

coupled with a slight increase of $T_{\rm r}$. There was virtually no change in Peak II. Conductivity increased as aging progressed but the hardness apparently decreased. The decrease in hardness is approximately equal to the scatter in the hardness measurements and may not be significant.

Figure 7 shows the result of aging at 120°C. After 6 h aging, Peak I is virtually non-existent. After 24 or 96 h, however, Peak I shows a large ΔH_r and a constant T_r of 192°C. The data in Table 2 shows that the ΔH_r after aging for 24 or 96 h was greater than in any of the previous aging treatments. This is consistent with the increased diffusivity of solute atoms at the higher aging temperature. The T_r for Peak I is also significantly higher than in any of the previous aging treatments. Peak II appears relatively sharp and displays a single maximum near 225°C after 6 h aging at 120°C. As aging proceeds, however, the shoulder on the high temperature side of the peak develops into a second peak. Note that the formation peaks with double maxima which were observed after the previously described lower temperature aging treatments had their second maximum located near 210°C, on the lower temperature side of the main 225°C peak, while the secondary maximum which develops after aging at 120°C is on the high temperature side, near 250°C. The maximum which develops near 250°C appears to be an additional formation reaction since the ΔC_{n} curve after 24 h aging lies below the 6 h curve. The hardness and conductivity both increase significantly during aging at 120°C.

3.3 150°C SECOND AGING TREATMENT

A second aging treatment of 8 h at 150°C was applied to several sets of the pre-aged samples. The first set of treatments consisted of 6, 24, and 96 h at 90°C followed by 8 h at 150°C. Results of DSC analysis of these three microstructures are shown in Fig. 8 and tabulated in Table 3. The results show that the microstructures were all quite similar, but ΔH_{Γ} and T_{Γ} for Peak I decreased slightly as the aging time at 90°C increased. Concurrently, ΔH_{Γ} for Peaks IIa and IIb increased and the hardness of the samples decreased slightly.

Similar trends were observed after aging for 6, 24, and 96 h at 120°C before aging 8 h at 150°C. The results in Fig. 9 and Table 3 show that the microstructures were similar but there might have been a slight decrease in

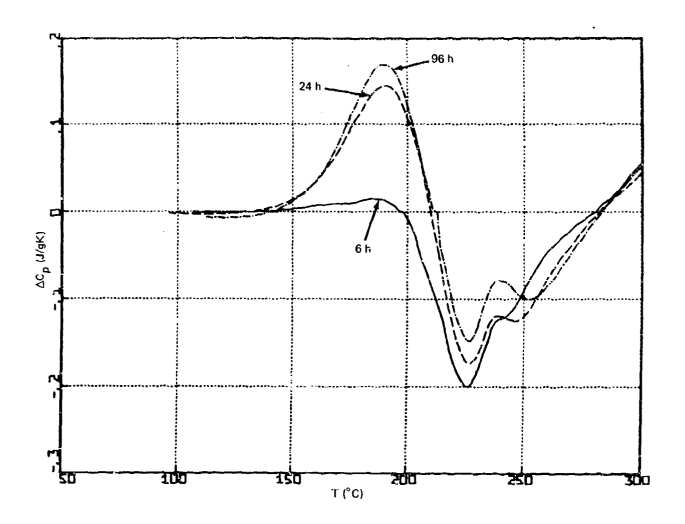


Fig. 7. DSC Traces From Solution Treated 7075 After Aging for 6, 24 and 96 Hours at 120° C.

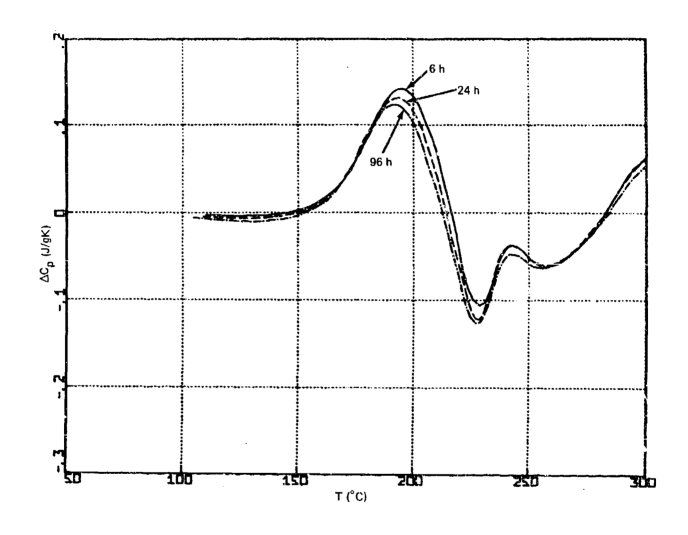


Fig. 8. DSC Traces From Solution Treated 7075 After Pre-Aging for 6, 24 and 96 Hours at 90°C and Final Aging for 8 Hours at 150°C.

Table 3. DSC Characteristics After Two Stage Aging Treatments Where Final Aging was at 150°C

Ag	Aging I Aging II				Peak I		Peak IIa		Peak 11b		
t (h)	T (°C)	t (h)	T (°C)	Hardness (R _B)	EC (% IACS)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)
6	90	8	150	86±1	33.8	196	4.8±.2	229	-1.8±.2	256	-1.6±.2
24	90	8	150	84±1	33.5	195	4.3±.3	229	-2.1±.3	257	-1.7±.1
96	90	8	150	83±2	33.7	193	4.1±.3	228	-2.4±.2	254	-1.8±.1
6	120	8	150	83±3	33.7	197±2	4.1±.3	228	-1.9±.2	258	-1.9±.2
24	120	8	150	83±1	33,5	195	4.2±.4	228	-2.1±.3	253±2	-2.1±.3
96	120	8	150	82±5	33.8	195	3.8±.4	228	-1.9±.2	253±2	-2,1±.2
6	75	8	150	82 ± 2	33.8	201	3.9±.1	230±2	-1.3±.3	260±3	-1.5±.2
6	85	8	150	83±4	33.5	194	3.7±.3	229	-2.5±.1	256	-2.3±.1
6	90	8	150	86±1	33.8	196	4.8±.2	229	-1.8±.2	256	-1.6±.2
6	95	8	150	84±2	33.4	194±2	3.5±.2	230	-2.3±.2	256	-2.3±.1
6	100	8	150	85±4	33.4	194	3.2±.3	228	-2.3±.3	254	-2,3±.1
6	105	8	150	80±1	33.9	196	4.7±.7	229	-1.7±.4	256	-1.6±.5
6	120	8	150	83±3	33.7	197±2	4.1±.3	228	-1.9±.2	256	-1.9±.2

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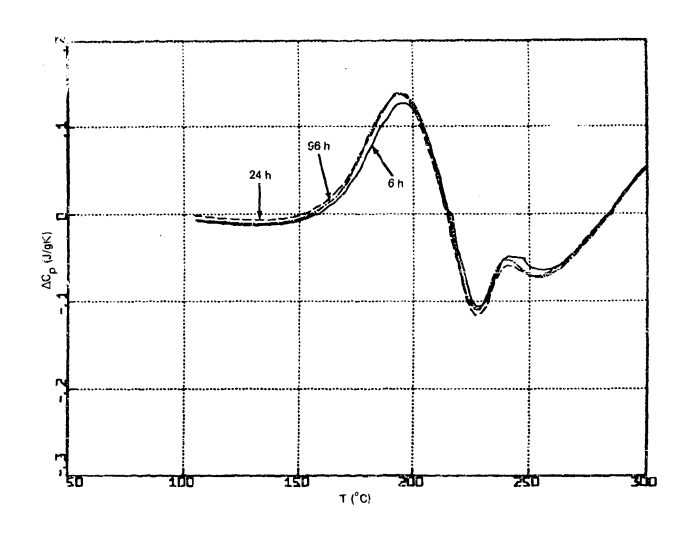


Fig. 9. DSC Traces From Solution Treated 7075 After Pre-Aging for 6, 24 and 96 Hours at 120°C and Final Aging for 8 Hours at 150°C.

 ΔH_Γ and T_Γ of Peak I. No significant change occurred in either Peak II or the hardness or the conductivity.

The third set of samples in this series were first aged for 6 h at 75, 85, 90, 95, 100, 105 and 120°C and then aged for 8 h at 150°C. Results from these samples are shown in Fig. 10 and tabulated in Table 3. The sample that was first aged at 75°C displayed a significantly smaller $^{\Delta}H_{\Gamma}$ for Peak II and a somewhat higher T_{Γ} for Peak I than the rest of the samples. For the remainder of the samples, the data do not seem to show any clear trends. There is some indication of a larger $^{\Delta}H_{\Gamma}$ for Peak I in the sample aged at 90 or 105°C.

3.4 COMMERCIAL TYPE AGING TREATMENT

This series of aging treatments consisted of small variations in aging temperatures centered around the values recommended by the Aluminum Association.

The first series of samples were aged for 4 h at 95°C and then 8 h at 150, 155 and 160°C. The DSC results are shown in Fig. 11 and tabulated in Table 4. The results show that increasing the second aging temperature to 155 or 160°C causes a significant increase in the $T_{\rm r}$ of Peak I and a very marked decrease in $\Delta H_{\rm r}$ of Peak II. The hardness and conductivity values also show this trend. Increasing the aging temperature to 155 or 160°C caused both the hardness and conductivity to increase compared to their value after aging at 150°C.

Figure 12 shows the results of aging for 4 h at 90, 95 and 100°C before aging for 8 h at 155°C. The curves in Fig. 12 indicate small differences between the microstructures, but the data in Table 4 indicate that there was no statistically significant difference between these three aging treatments. Similarly, the hardness and conductivity values do not show any significant changes.

The results from the final batch of samples in this group are shown in Fig. 13. These samples were aged for 8 h at 95, 100, 105 and 110°C before aging for 24 h at 165°C. No significant difference was observed in the DSC results or in the electrical conductivity or hardness of these samples.

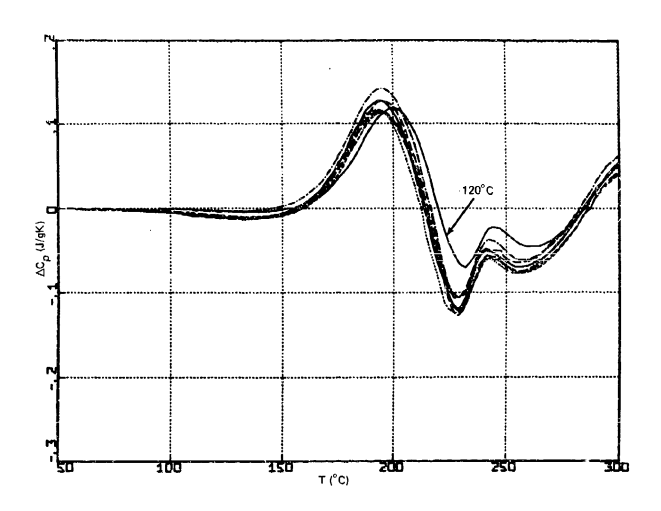


Fig. 10. DSC Traces From Solution Treated 7075 After Pre-Aging for 6 Hours at 75, 85, 90, 95, 100, 105 and 120°C and Final Aging for 8 Hours at 150°C. Most of the Curves are Very Similar and Thus Could not be Labeled.

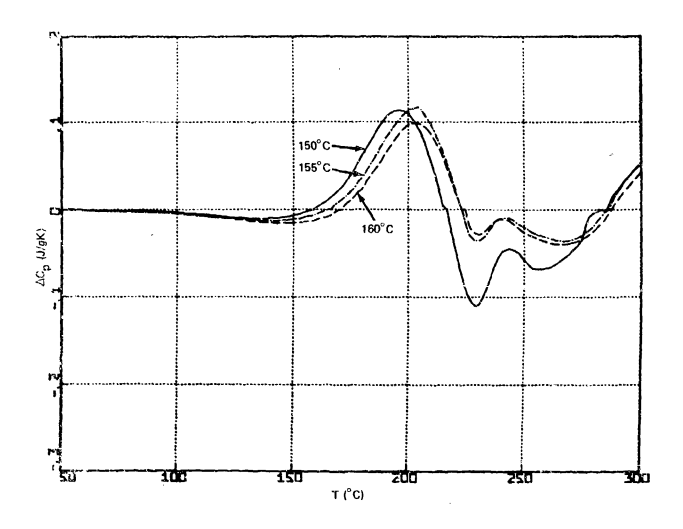


Fig. 11. DSC Traces From Solution Treated 7075 After Pre-Aging for 4 Hours at 95°C and Final Aging for 8 Hours at 150, 155 and 160°C.

Table 4. DSC Characteristics After Commercial Type Aging Treatments

Agi	Aging I Aging II				Peak I		Peak IIa		Peak IIb		
t (h)	T (°C)	t (h)	T (°C)	Hardness (R _B)	EC (% IACS)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)	T _r (°C)	ΔH _r (J/g)
4	95	8	150	79±3	33.5	195±1	3.5±.1	229	-2.1±.3	255	-2.1±.2
4	95	8	155	83±4	36.0	205	3.3±.5	230	-,2±.2	263	9±.5
4	95	8	160	84±3	35.6	203±2	3.6±.1	230±2	5±.1	266±2	-1.1±.1
4	90	8	155	84±2	36.5	205±2	3.3±.5	231±2	1±.2	267±3	6±.3
4	95	8	155	83±4	36.0	205	3.3±.5	230	2±.2	263	9±.5
4	100	8	155	81±4	35.9	202±2	3.5±.5	231	.4±.1	266±3	-1.0±.1
8	95	24	165	82±2	38.6	20€	4,2±.2				
8	165	24	165	79±4	38.6	208	3.9±.1				
8	105	24	165	80±2	39.0	208	4.0±.5				
8	i 10	24	165	83±3	38.7	209	4.1±.1				
RRA	4	SURI	ACE	84±2		205	6.5±.1				
RRA	Α	MID- THIC	KNESS	87±1		210	6.2±.2				

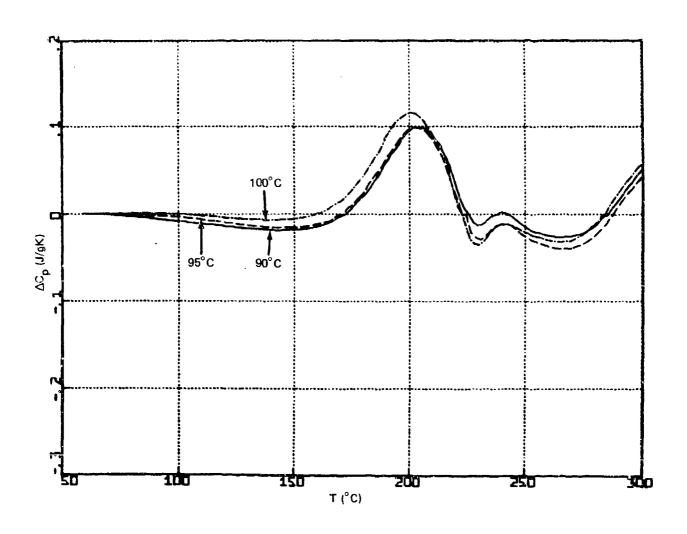


Fig. 12. DSC Traces From Solution Treated 7075 After Pre-Aging for 4 Hours at 90, 95 and 100°C and Final Aging for 8 Hours at 155°C.

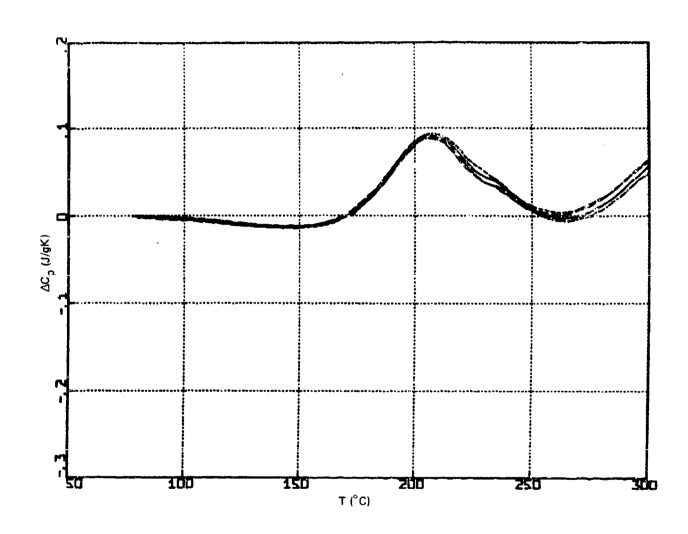


Fig. 13. DSC Traces From Solution Treated 7075 After Pre-Aging for 8 Hours at 95, 100, 105 and 110°C and Final Aging for 24 Hours at 165°C.

3.5 RETROGRESSION AND REAGING TREATMENT

Samples were taken from near the surface and at the mid-thickness of a 1 in. thick piece of 7075-T651 plate that had been RRA treated by Lockheed-California Company. The DSC results from these samples are shown in Fig. 14 and consist solely of a dissolution peak at approximately 205°C. There was little difference between the surface and the center of the sample. The shape of the DSC trace is similar to the T7 type of microstructure shown in Fig. 13, but the $H_{\rm T}$ of Peak I is significantly greater in the RRA samples.

3.6 TRANSMISSION ELECTRON MICROSCOPY RESULTS

The results of the TEM studies of grain boundary precipitates are listed in Table 5. The samples investigated had received the following aging treatments: a T6 treatment (24 h at 120°C), a two stage T6 treatment (4 h at 95°C and 8 h at 155°C), a T73 treatment (8 h at 105°C and 24 h at 165°C), and a laboratory RRA treatment of 24 h at 120°C plus 60 s at 240°C plus 24 h at 120°C. Note that this may not be the optimum RRA treatment. The data show large variations in the numbers and sizes of the precipitates between different boundaries in the same thin foil and between different thin foils from the same sample. Thus, the standard deviations listed in Table 5 are so large that it is difficult to draw conclusions. There is some indication that the T7 type of heat treatment resulted in the largest average grain boundary precipitate diameter, the largest area fraction covered and the smallest number of particles per unit area. The standard T6 aging and the RRA type of heat treatment both appeared to give similar results of smallest particle size, smallest fraction coverage and largest particle density. The two stage T6 type of aging treatment resulted in grain boundary precipitate characteristics which were intermediate between the two extremes.

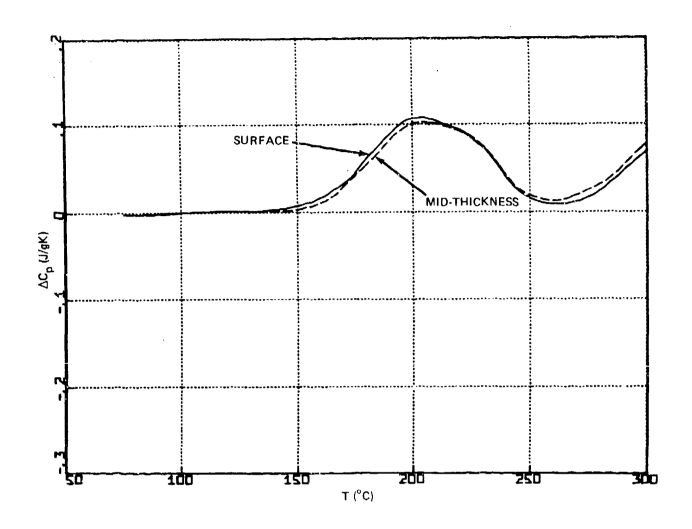


Fig. 14. DSC Traces From the Surface and Mid-Thickness of a One Inch Thick Piece of 7075-T651 that Received the Patented Retrogression and Re-Aging (RRA) Treatment.

Table 5. Characteristics of the Grain Boundary Precipitates

Temper	N	N _A (μm²) ⁻¹	₹ (nm)	Af
Т6	635	302±190	28±13	0.16±.06
Two Stage T6	401	130±80	55±19	0.23±.09
RRA*	545	372±185	34±16	0.21±.05
T73	102	80±50	66±20	0.29±.03

^{*}Laboratory treated T6 material that was given a retrogression treatment if 60 s at 240°C and then re-aged for 24 h at 120°C.

4. DISCUSSION

The results of this program contain a great deal of information about the microstructures produced during aging of 7075. In particular, the effects of the various low temperature pre-aging treatments show a rather complex pattern of microstructural evolution. Previously published DSC studies of 7075 in which the DSC signatures were compared to the TEM microstructures dealt only with aging temperatures of 120°C or higher (Ref 5). Thus, there is no firm basis upon which to assign specific reactions to specific reaction peaks in the data. Numerous observations suggest themselves, but it would probably be misleading to speculate on the nature of these low temperature microstructures without further information. An example of the difficulty of interpreting the low temperature aging results can be found in Fig. 2 where aging for a given time at increasing temperatures results in a smaller and smaller dissolution enthalpy for Peak I. This does not necessarily indicate that less precipitate is formed at the higher temperature, but may simply show that the dissolution and formation peaks are overlapping. Without detailed knowledge of the reactions involved, further comment is not justified. The results of the two-stage aging treatments, however, show several significant trends.

The results from experiments in which pre-aging was carried out for 6, 24 and 96 h at 90 or 120°C before final aging for 8 h at 150°C show that the final microstructures were quite similar for all of these conditions. However, the dissolution enthalpy for Peak I, the strengthening precipitate, is a maximum for the shortest pre-age at the lower temperature. This dissolution enthalpy decreases regularly as the time and temperature of preaging increase. The hardness values lead to the same general conclusion, with the shortest pre-age at the lowest temperature giving the hardest structure. The microstructure produced by 6 h at 90°C plus 8 h at 150°C is very similar to the standard T6 (24 h at 120°C) both in hardness and in DSC characteristics with the exception of Peak II which was smaller in the two-stage sample. These microstructural results can be understood on the basis of the Pashley et al model for two stage aging (Ref 15) since a finer

low temperature microstructure will result in a finer final microstructure. The Lorimer and Nicholson model (Ref 14) would predict the opposite behavior, since a longer preaging treatment should result in a larger fraction of G.P. zones which can survive the up-quench and hence nucleate a finer final microstructure, which would presumably be stronger.

The results of the two-stage aging experiments in which pre-aging was carried out for 6 h at various temperatures before final aging at 150°C again show similar final microstructures, but with some trends suggested by the data. The lowest aging temperature, 75°C, when compared to the standard T6 (24 h at 120°C) appears to result in a slightly overaged, weaker microstructure since Tr for Peak I is above 200°C and AHr for Peak II is significantly reduced. As the pre-aging temperature increases, the characteristics of the samples improve and then oscillate. This behavior, if it is genuine and not simply due to inaccuracies and error accumulation, can be explained on the basis of inadequate pre-aging at 75°C and, subsequently, the formation of different precipitate phases at different preaging temperatures. A change from G.P. zone microstructures to n" microstructures in this temperature range has been documented for medium strength Al-Zn-Mg alloys (Ref 23 and 24) and suggested for 7075 (Ref 11). Clarification of the nature of the microstructures produced during the pre-aging treatments and an accurate placement of the phase boundaries would aid in understanding this result.

The results of varying the final aging temperature from 150 to $160\,^{\circ}\text{C}$ show significant microstructural effects in this temperature range as shown in Fig. 11. Changing the final aging temperature from 150 to $155\,^{\circ}\text{C}$ resulted in an approximately four fold decrease in the area of Peak II with only a small effect on Peak I. This result implies that the rate of formation of η and/or η is significantly greater at $155\,^{\circ}\text{C}$ than at $150\,^{\circ}\text{C}$. Increasing the temperature further to $160\,^{\circ}\text{C}$ had little additional effect. Pre-aging for 4 h at 90 to $100\,^{\circ}\text{C}$ with final aging of 8 h at $155\,^{\circ}\text{C}$ is specified in the Aluminum Standards and Data publication (Ref 4) as an alternate means of producing the T6 temper. This result shows that increasing the final aging temperature from $150\,^{\circ}\text{C}$ as suggested by DiRusso et al (Ref 1) to $155\,^{\circ}\text{C}$ as recommended by the Aluminum Association (Ref 4) has a pronounced effect on the final microstructure. The effect of varying the pre-aging temperature between 90

and 100°C was not as pronounced, as can be seen in Fig. 12. The differences indicated in Fig. 12 are not supported by the actual enthalpies listed in Table 4. These results agree with the tensile data presented by Buratti et al (Ref 3). They showed that variations in the pre-aging treatment did not have much of effect on the final properties but that changing the final aging temperature caused significant shifts in the aging curves. This same observation holds true for the two-stage T73 (8 h at 105°C plus 24 h at 165°C) and T76 (8 h at 100°C plus 24 h at 165°C) type treatments shown in Fig. 13. No significant differences were observed between the characteristics of the microstructures produced by varying the pre-aging temperature between 95 and 110°C when the final aging treatment was 24 h at 165°C.

The DSC characteristics of the microstructures produced by the patented RRA treatment appear to be similar to those of the T7 type treatments. However, the dissolution enthalpy for Peak I in the RRA material is approximately 50% greater and the peak is much broader. We have observed previously in this work, and in other studies (Ref 6), that the hardness is related to the enthalpy of Peak I, thus the larger dissolution enthalpy in the RRA material is expected to result in greater hardness, as observed. These results indicate that the RRA treatment results in η and η microstructures which are more similar to T7 microstructures than to T6, but which are harder than the η and η microstructures produced by two stage, T7 type treatments.

The differences between the various commercially used microstructures are shown in Fig. 15. The most pronounced difference is thought to be in Peak II, the formation peak. For the T7 and RRA treatments, Peak II is absent. This indicates that these microstructures consist entirely of η' and η . The two stage T6 shows a small Peak II, while the standard T6 shows a large Peak II. Thus, while the T7 and RRA microstructures are exclusively η' and η , the two stage T6 contains less of these phases and the ordinary T6 least of all. In addition, Peak I is different in the various conditions. When compared to the ordinary T6, Peak I in the two stage T6 is broader and shifted to a higher temperature. This is indicative of a larger average particle size and/or a greater proportion of η' in the two stage microstructure. Peak I for the T7 microstructures is somewhat broader as is typical of the η' usually observed in this temper (Ref 5). This indicates a broader particle size distribution

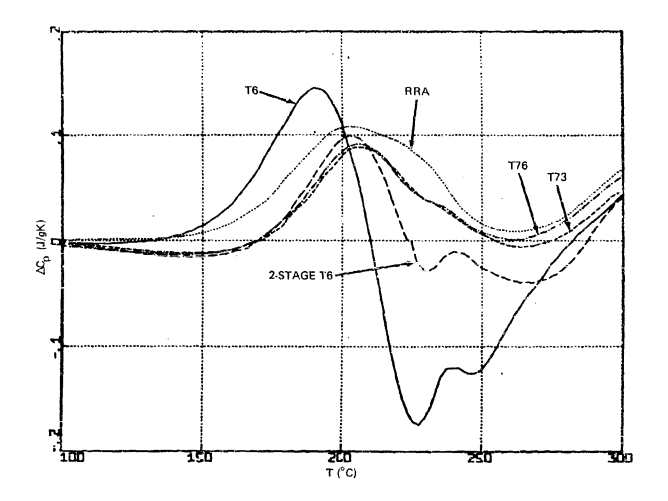


Fig. 15. DSC Tracas From Solution Treated 7075 that Received a T6 Aging Treatment (24 Hours at 120°C), a Two-Stage T6 Treatment (4 Hours at 95°C and 8 Hours at 155°C), a T76 Treatment (8 Hours at 100°C and 24 Hours at 165°C) and a T73 Treatment (8 Hours at 105°C and 24 Hours at 165°C) and From a Sample of 7075-T651 Which Received the Patented RRA Treatment.

or may simply be the result of the absence of overlap with Peak II. Peak I in the RRA microstructure is broader still and appears to be a superposition of multiple dissolution peaks, suggesting that the RRA microstructure consists of a mixture of several precipitate phases. All of the tempers are mixtures of precipitate phases to some extent, but the RRA appears to contain the greatest variety.

In summary therefore, the high final aging temperature of the two stage T7 results in a predominately n' microstructure which is more coarsely distributed then those microstructures produced at lower temperatures. The two-stage T6, which receives a final aging treatment at 155°C, succeeds in conserving the relatively fine microstructure it inherits from the low temperature pre-age. The ordinary T6 contains the finest distribution of strengthening precipitates, predominately G.P. zones, but it has the least amount of more stable η' and η phases. The RRA treatment appears to dissolve the G.P. zones inherited from the T6 starting microstructure, but, since the final aging treatment is performed at 120°C, excessive coarsening of the η' and η components of the microstructure does not occur and the final result is both fine scale and stable.

The relationship of these results to the stress corrosion susceptibility of the rarious tempers will now be considered. Unfortunately, the stress corrosion data for these samples is not yet available, but, for the purposes of this short discussion we will assume on the basis of previously published results (Ref 1, 2, and 29) that the T7 and the RRA tempers are the most stress corrosion resistant, that the ordinary T6 is the least resistant and the two stage T6 is intermediate. Various authors have found that the stress corrosion susceptibility of Ai-Zn-Mg alloys is controlled by the characteristics of the grain boundary precipitates (Ref 30 through 35), the matrix microstructure (Ref 36), or by factors related to hydrogen embrittlement (Ref 37 through 41).

If grain boundary precipitate density and size were the primary factor, then the data in Table 5 would lead us to expect best stress corrosion resistance for the T73, worst for the T6 and RRA, and intermediate resistance for the two stage T6. This is not in agreement with the previously published results for these conditions mentioned above. If matrix microstructure were of

greatest importance, then we would predict a different result. A uming that the DSC characteristics of the T7 microstructures are the desirable characteristics for stress corrosion resistance and the DSC characteristics of the T6 microstructure are the undesirable ones, then we would predict that the RRA should have a stress corrosion resistance equivalent to that of the T7. This prediction is based on the enthalpy of Peak II, which is absent in the T7 and RRA microstructures and which is maximum for the T6. The two stage T6 would be expected to have intermediate stress corrosion resistance on this basis. This prediction agrees with the expected ranking mentioned above.

Recent work on stress corrosion has emphasized the role of hydrogen, in particular, the effects of magnesium segregated to grain boundaries (Ref. 41 and 42) and the role of grain boundary precipitates (Ref 40). If magnesium segregation is of primary importance, and if it is assumed that the segregation is of the non-equilibrium, vacancy aided type which occurs during quenching (Ref 43 and 44), then aging at elevated temperatures will allow re-establishment of equilibrium solute partitioning by diffusion. Rough calculations of the root mean square diffusion distance of magnesium during the aging treatments indicate that the T7 allows the greatest diffusion distance, the RRA allows approximately 80% as much, the two-stage 16 allow-35% as much, and the T6 only allows 10% as much. Thus, this theory correctly predicts the expected ranking. Finally, Christodoulou and Flower (Ref 40) have found that grain boundary n particles larger than 20 nm are effective as nucleation sites for hydrogen bubbles, and thus decrease the amount of hydrogen dissolved in the matrix and grain boundaries thereby reducing the stress corrosion susceptibility. Our characterization of the grain boundary precipitates was not quantitative enough to determine the relative fraction of precipitates greater than 20 nm, thus our data do not allow comparison to this theory.

5. SUMMARY

- 1. For two-stage aging treatments in which the final aging is for 8 h at 150°C, pre-aging for 6 h at 90°C produces a slightly finer and harder microstructure than longer treatments at 90°C or equivalent treatments at 120°C. This is in agreement with the Pashley et al model of two-stage aging.
- 2. For two-stage treatments where the first aging treatment is 4 h at 95°C, increasing the final aging temperature from 150 to 155 or 160°C results in a significant increase in the amount of η' in the microstructure. Variations in the pre-aging treatment had much less of an effect.
- 3. The RRA heat treaqtment converts the T6, predominately G.P. zone microstructure, to a more stable T7 type of microstructure; but the RRA microstructure consists of smaller precipitates and is more heterogeneous than the T7.
- 4. The major differences between the T6, two-stage T6, RRA and T73 microstructures appear to be in the relative proportions of G.P. zones, n' and n. The percentage of G.P. zones is greatest in T6, least in T73 and RRA and intermediate in the two-stage T6.
- 5. The stress corrosion susceptibility of 7075 appears to be more influenced by the matrix microstructure than by the grain boundary precipitate size and spacing.

RECOMMENDATIONS FOR FUTURE WORK

The observations made in the course of this work have raised several technical issues which should be addressed in order to complement the experiments already performed and to maximize the scientific benefit of this effort. They are:

- 1. Comparison stress corrosion testing of the four commercial tempers, (T6, two-stage T6, T73 and RRA) should be performed. The results of these stress corrosion tests are necessary for interpretation of the DSC results obtained in this work. The two-stage T6 should not be overlooked in the stress corrosion testing. As shown in the discussion, stress corrosion data from the two-stage T6 condition is a key factor in the separation of the various microscructural effects that have been observed, and would be of great utility in determining which particular microstructural characteristic is of greatest significance in the stress corrosion behavior of 7075 in a high strength condition.
- 2. Additional DSC studies of the microstructures in the various stages of the RRA treatment in 7075 should be performed and correlated with stress corrosion behavior. As shown in our progress report, DSC is a very rapid and sensitive indicator of the extent of retrogression and, as such, it can be of great use in establishing the correct processing condtions for a particular part. Once the optimum microstructural conditions after retrogression and after reaging are established, these data could be used for rapid and precise determination of processing parameters for parts of different thicknesses, geometries or prior microstructural conditions.
- 3. DSC studies of the various stages of RRA treatments in 7050 should be performed and correlated with stress corrosion testing. Since the T6 microstructure of 7050 has a DSC signature which closely resembles that of the two-stage T6 in 7075, it is expected that RRA may provide less improvement in the stress corrosion behavior of 7050 than of

- 7075. Data from these tests would allow generalization of the conclusions about the relationship between microstructure and stress corrosion susceptibility in 7075 to the 7000 series.
- 4. Additional TEM studies should be performed to establish the precise identity of the precipitates formed during low temperature aging of 7075, and these data should be correlated with the DSC results in order to allow a better understanding of the structures obtained in the pre-aging treatments and to enhance the utility of the DSC technique.
- 5. Quantitative transmission electron microscopy studies using dark field techniques should be performed in samples representative of the four commercial type aging treatments. The objectives of this work would be to quantitatively determine the number density and size distribution of the G.P. zones, η'' , η' and η precipitates both in the matrix and grain boundaries of the four microstructures. The results of this study when compared with stress corrosion and mechanical property data will establish the precise microstructural conditions which lead to best properties and should allow even further microstructural optimization.

7. REFERRENCES

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